Hybrid MPI and OpenMP
Parallel Programming

MPI + OpenMP and other models
on clusters of SMP nodes

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“Hochleistungsrechnen“
Prof. Dr. habil Thomas Ludwig, Deutsches Klimarechenzentrum (DKRZ),
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Outline

• Introduction / Motivation 2
• Programming models on clusters of SMP nodes 6
• Case Studies / Benchmark results 13
• Mismatch Problems 23
• Opportunities: Application categories that can benefit from hybrid parallelization 52
• Thread-safety quality of MPI libraries 62
• Other options on clusters of SMP nodes 67
• Summary 81
• Appendix 89
Motivation

- Efficient programming of clusters of SMP nodes
  - SMP nodes:
    - Dual/multi core CPUs
    - Multi CPU shared memory
    - Multi CPU ccNUMA
    - Any mixture with shared memory programming model
- Hardware range
  - mini-cluster with dual-core CPUs
  - ...
  - large constellations with large SMP nodes
    - ... with several sockets (CPUs) per SMP node
    - ... with several cores per socket
  - Hierarchical system layout
- Hybrid MPI/OpenMP programming seems natural
  - MPI between the nodes
  - OpenMP inside of each SMP node
Motivation

- Which programming model is fastest?
- MPI everywhere?
- Fully hybrid MPI & OpenMP?
- Something between? (Mixed model)
- Often hybrid programming slower than pure MPI
  - Examples, Reasons, …
Goals of this tutorial

- Sensitize to problems on clusters of SMP nodes
  - see sections → Case studies
  - → Mismatch problems
- Technical aspects of hybrid programming
  - see sections → Programming models on clusters
  - → Examples on hybrid programming
- Opportunities with hybrid programming
  - see section → Opportunities: Application categories that can benefit from hybrid paralleliz.
- Issues and their Solutions
  - with sections → Thread-safety quality of MPI libraries
  - → Tools for debugging and profiling for MPI+OpenMP

• Less frustration & • More success with your parallel program on clusters of SMP nodes
Outline

- Introduction / Motivation
- Programming models on clusters of SMP nodes
  - Case Studies / Benchmark results
  - Mismatch Problems
  - Opportunities:
    Application categories that can benefit from hybrid parallelization
  - Thread-safety quality of MPI libraries
  - Other options on clusters of SMP nodes
  - Summary
Major Programming models on hybrid systems

- Pure MPI (one MPI process on each core)
- Hybrid MPI+OpenMP
  - shared memory OpenMP
  - distributed memory MPI
- Other: Virtual shared memory systems, PGAS, HPF, ...
- Often **hybrid programming (MPI+OpenMP)** slower than **pure MPI**
  - why?

**MPI**
- Sequential program on each core
- Explicit **Message Passing** by calling **MPI_Send & MPI_Recv**
- local data in each process

**OpenMP** (shared data)
- some_serial_code
- #pragma omp parallel for
- for (j=...;...; j++)
- block_to_be_parallelized
- again_some_serial_code
- by calling **MPI_Send & MPI_Recv**
- local data in each process

**Node Interconnect**
- Master thread, other threads
- • • • sleeping • • •
- MPI between the nodes via node interconnect
- OpenMP inside of the SMP nodes

Hybrid Parallel Programming
Rolf Rabenseifner
Slide 7 / 88
Parallel Programming Models on Hybrid Platforms

- **pure MPI**
  - one MPI process on each core

- **hybrid MPI+OpenMP**
  - MPI: inter-node communication
  - OpenMP: inside of each SMP node

- **OpenMP only**
  - distributed virtual shared memory

No overlap of Comm. + Comp.
- MPI only outside of parallel regions of the numerical application code

Overlapping Comm. + Comp.
- MPI communication by one or a few threads while other threads are computing

Masteronly
- MPI only outside of parallel regions
Pure MPI

Advantages
- No modifications on existing MPI codes
- MPI library need not to support multiple threads

Major problems
- Does MPI library uses internally different protocols?
  - Shared memory inside of the SMP nodes
  - Network communication between the nodes
- Does application topology fit on hardware topology?
- Unnecessary MPI-communication inside of SMP nodes!

Discussed in detail later on in the section Mismatch Problems
Hybrid Masteronly

Advantages
- No message passing inside of the SMP nodes
- No topology problem

for (iteration ....)
{
    #pragma omp parallel
    numerical code
    /*end omp parallel */

    /* on master thread only */
    MPI_Send (original data to halo areas in other SMP nodes)
    MPI_Recv (halo data from the neighbors)
} /*end for loop

Major Problems
- All other threads are sleeping while master thread communicates!
- Which inter-node bandwidth?
- MPI-lib must support at least MPI_THREAD_FUNNELED

→ Section Thread-safety quality of MPI libraries
Overlapping Communication and Computation

MPI communication by one or a few threads while other threads are computing

if (my_thread_rank < ...) {
    MPI_Send/Recv....
    i.e., communicate all halo data
} else {
    Execute those parts of the application
    that do not need halo data
    (on non-communicating threads)
}

Execute those parts of the application
that need halo data
(on all threads)
Pure OpenMP (on the cluster)

- Distributed shared virtual memory system needed
- Must support clusters of SMP nodes
- e.g., Intel® Cluster OpenMP
  - Shared memory parallel inside of SMP nodes
  - Communication of modified parts of pages at OpenMP flush (part of each OpenMP barrier)

i.e., the OpenMP memory and parallelization model is prepared for clusters!

Experience: Mismatch section
Outline

- Introduction / Motivation
- Programming models on clusters of SMP nodes

**Case Studies / Benchmark results**
- The Multi-Zone NAS Parallel Benchmarks
  - Gabriele Jost (University of Texas, TACC/Naval Postgraduate School, Monterey CA)
- Micro Benchmarks
  - Georg Hager (Regionales Rechenzentrum Erlangen, RRZE)

- Mismatch Problems
- Opportunities: Application categories that can benefit from hybrid parallel
- Thread-safety quality of MPI libraries
- Other options on clusters of SMP nodes
- Summary
The Multi-Zone NAS Parallel Benchmarks

- Aggregate sizes:
  - Class D: 1632 x 1216 x 34 grid points
  - Class E: 4224 x 3456 x 92 grid points

- **BT-MZ:** *(Block tridiagonal simulated CFD application)*
  - Alternative Directions Implicit (ADI) method
  - #Zones: 1024 (D), 4096 (E)
  - Size of the zones varies widely:
    - large/small about 20
    - requires multi-level parallelism to achieve a good load-balance

- **LU-MZ:** *(LU decomposition simulated CFD application)*
  - SSOR method (2D pipelined method)
  - #Zones: 16 (all Classes)
  - Size of the zones identical:
    - no load-balancing required
    - limited parallelism on outer level

- **SP-MZ:** *(Scalar Pentadiagonal simulated CFD application)*
  - #Zones: 1024 (D), 4096 (E)
  - Size of zones identical
    - no load-balancing required

Expectations:

- Pure MPI: Load-balancing problems!
- Good candidate for MPI+OpenMP
- Limited MPI Parallelism: \( \rightarrow \) MPI+OpenMP increases Parallelism
- Load-balanced on MPI level: Pure MPI should perform best
Sun Constellation Cluster Ranger (1)

- Located at the Texas Advanced Computing Center (TACC), University of Texas at Austin (http://www.tacc.utexas.edu)
- 3936 Sun Blades, 4 AMD Quad-core 64bit 2.3GHz processors per node (blade), 62976 cores total
- 123TB aggregate memory
- Peak Performance 579 Tflops
- InfiniBand Switch interconnect
- Sun Blade x6420 Compute Node:
  - 4 Sockets per node
  - 4 cores per socket
  - HyperTransport System Bus
  - 32GB memory
NPB-MZ Class E Scalability on Sun Constellation

- Scalability in Mflops
- MPI/OpenMP outperforms pure MPI
- Use of numactl essential to achieve scalability

**SP**
- Pure MPI is already load-balanced.
- But hybrid 9.6% faster, due to smaller message rate at NIC
- Cannot be build for 8192 processes!

**BT**
- Significant improvement (235%):
  - Load-balancing issues solved with MPI+OpenMP

**Hybrid:**
- **SP:** still scales
- **BT:** does not scale

Slides, courtesy of Gabriele Jost, TACC, Austin, USA
NUMA Control: Process Placement

- Affinity and Policy can be changed externally through `numactl` at the socket and core level.

**Command:** `numactl <options> ./a.out`

**Example:**
- `numactl -N 1 ./a.out`
- `numactl --c 0,1 ./a.out`
NUMA Operations: Memory Placement

Memory allocation:
- MPI
  - local allocation is best
- OpenMP
  - Interleave best for large, completely shared arrays that are randomly accessed by different threads
  - local best for private arrays
- Once allocated, a memory-structure is fixed

Example: `numactl -N 1 -l ./a.out`
### NUMA Operations (cont. 3)

<table>
<thead>
<tr>
<th>Operation</th>
<th>Command</th>
<th>Option</th>
<th>Arguments</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Socket Affinity</td>
<td>numactl</td>
<td>-N</td>
<td>{0,1,2,3}</td>
<td>Only execute process on cores of this (these) socket(s).</td>
</tr>
<tr>
<td>Memory Policy</td>
<td>numactl</td>
<td>-l</td>
<td>{no argument}</td>
<td>Allocate on current socket.</td>
</tr>
<tr>
<td>Memory Policy</td>
<td>numactl</td>
<td>-i</td>
<td>{0,1,2,3}</td>
<td>Allocate round robin (interleave) on these sockets.</td>
</tr>
<tr>
<td>Memory Policy</td>
<td>numactl</td>
<td>--preferred=</td>
<td>{0,1,2,3}</td>
<td>Allocate on this socket; fallback to any other if full.</td>
</tr>
<tr>
<td>Memory Policy</td>
<td>numactl</td>
<td>-m</td>
<td>{0,1,2,3}</td>
<td>Only allocate on this (these) socket(s).</td>
</tr>
<tr>
<td>Core Affinity</td>
<td>numactl</td>
<td>-C</td>
<td>{0,1,2,3, 4,5,6,7, 8,9,10,11, 12,13,14,15}</td>
<td>Only execute process on this (these) Core(s).</td>
</tr>
</tbody>
</table>
Intra-node MPI characteristics: IMB Ping-Pong benchmark

- Code (to be run on 2 processors):

```fortran
wc = MPI_WTIME()
do i=1,NREPEAT
    if(rank.eq.0) then
        MPI_SEND(buffer,N,MPI_BYTE,1,0,MPI_COMM_WORLD,ierr)
        MPI_RECV(buffer,N,MPI_BYTE,1,0,MPI_COMM_WORLD, &
                  status,ierr)
    else
        MPI_RECV(…)
        MPI_SEND(…)
    endif
endo
wc = MPI_WTIME() - wc
```

- Intranode (1S): `mpirun -np 2 -pin "1 3" ./a.out`
- Intranode (2S): `mpirun -np 2 -pin "2 3" ./a.out`
- Internode: `mpirun -np 2 -pernode ./a.out`
IMB Ping-Pong: Latency
Intra-node vs. Inter-node on Woodcrest DDR-IB cluster (Intel MPI 3.1)

Affinity matters!

Slides, courtesy of Georg Hager, RRZE, Erlangen
IMB Ping-Pong: Bandwidth Characteristics

Intra-node vs. Inter-node on Woodcrest DDR-IB cluster (Intel MPI 3.1)

- **between two cores of one socket**
- **between two sockets of one node**
- **between two nodes via InfiniBand**

**Affinity matters!**

Slides, courtesy of Georg Hager, RRZE, Erlangen
Outline

• Introduction / Motivation
• Programming models on clusters of SMP nodes
• Case Studies / Benchmark results

• **Mismatch Problems**

• Opportunities:
  Application categories that can benefit from hybrid parallelization
• Thread-safety quality of MPI libraries
• Other options on clusters of SMP nodes
• Summary
Mismatch Problems

- None of the programming models fits to the hierarchical hardware (cluster of SMP nodes)
- Several mismatch problems → following slides
- Benefit through hybrid programming → Opportunities, see next section
- Quantitative implications → depends on your application

<table>
<thead>
<tr>
<th>Examples:</th>
<th>No.1</th>
<th>No.2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benefit through hybrid (see next section)</td>
<td>30%</td>
<td>10%</td>
</tr>
<tr>
<td>Loss by mismatch problems</td>
<td>-10%</td>
<td>-25%</td>
</tr>
<tr>
<td>Total</td>
<td>+20%</td>
<td>-15%</td>
</tr>
</tbody>
</table>

In most cases: Both categories!
The Topology Problem with pure MPI

one MPI process on each core

Application example on 80 cores:
- Cartesian application with $5 \times 16 = 80$ sub-domains
- On system with $10 \times$ dual socket $\times$ quad-core

- 17 x inter-node connections per node
- 1 x inter-socket connection per node

Sequential ranking of MPI_COMM_WORLD

Does it matter?
The Topology Problem with pure MPI
one MPI process on each core

Application example on 80 cores:
- Cartesian application with 5 x 16 = 80 sub-domains
- On system with 10 x dual socket x quad-core

Never trust the default !!!

32 x inter-node connections per node
0 x inter-socket connection per node

Round robin ranking of MPI_COMM_WORLD
The Topology Problem with pure MPI

Application example on 80 cores:
- Cartesian application with $5 \times 16 = 80$ sub-domains
- On system with $10 \times$ dual socket $\times$ quad-core

- 12 x inter-node connections per node
- 4 x inter-socket connection per node

Bad affinity of cores to thread ranks
The Topology Problem with pure MPI
one MPI process on each core

Application example on 80 cores:
- Cartesian application with $5 \times 16 = 80$ sub-domains
- On system with $10 \times$ dual socket $\times$ quad-core

Two levels of domain decomposition

12 x inter-node connections per node
2 x inter-socket connection per node

Good affinity of cores to thread ranks
The Topology Problem with

Problem

– Does application topology inside of SMP parallelization fit on inner hardware topology of each SMP node?

Solutions:

– Domain decomposition inside of each thread-parallel MPI process, and
– first touch strategy with OpenMP

Successful examples:

– Multi-Zone NAS Parallel Benchmarks (MZ-NPB)
The Topology Problem with hybrid MPI+OpenMP

Application example:
- Same Cartesian application aspect ratio: 5 x 16
- On system with 10 x dual socket x quad-core
- 2 x 5 domain decomposition

MPI: inter-node communication
OpenMP: inside of each SMP node

+ 3 x inter-node connections per node, but ~ 4 x more traffic
+ 2 x inter-socket connection per node

Affinity of cores to thread ranks !!!
Inside of an SMP node

- 2nd level of domain decomposition: OpenMP
- 3rd level: 2nd level cache
- 4th level: 1st level cache

Optimizing the numerical performance
The Mapping Problem with mixed model

Several multi-threaded MPI process per SMP node:

Problem
- Where are your processes and threads really located?

Solutions:
- Depends on your platform,
- e.g., with `numactl`
  
  → Case study on Sun Constellation Cluster Ranger with BT-MZ and SP-MZ

Further questions:
- Where is the NIC\(^1\) located?
- Which cores share caches?

\(^1\) NIC = Network Interface Card
Unnecessary intra-node communication

Problem:
- If several MPI processes on each SMP node
  → unnecessary intra-node communication

Solution:
- Only one MPI process per SMP node

Remarks:
- MPI library must use appropriate fabrics/protocol for intra-node communication
- Intra-node bandwidth higher than inter-node bandwidth
  → problem may be small
- MPI implementation may cause unnecessary data copying
  → waste of memory bandwidth
Sleeping threads and network saturation

Problem 1:
- Can the master thread saturate the network?
  Solution:
  - If not, use mixed model
  - i.e., several MPI processes per SMP node

Problem 2:
- Sleeping threads are wasting CPU time
  Solution:
  - Overlapping of computation and communication

Problem 1 & 2 together:
- Producing more idle time through lousy bandwidth of master thread

```c
for (iteration ....) {
    #pragma omp parallel
    numerical code
    /*end omp parallel */

    /* on master thread only */
    MPI_Send (original data to halo areas in other SMP nodes)
    MPI_Recv (halo data from the neighbors)
} /*end for loop*/
```
OpenMP: Additional Overhead & Pitfalls

- Using OpenMP
  - may prohibit compiler optimization
  - may cause significant loss of computational performance
- Thread fork / join overhead
- On ccNUMA SMP nodes:
  - Loss of performance due to missing memory page locality or missing first touch strategy
  - E.g. with the masteronly scheme:
    - One thread produces data
    - Master thread sends the data with MPI
    - data may be internally communicated from one memory to the other one
- Amdahl’s law for each level of parallelism
- Using MPI-parallel application libraries? → Are they prepared for hybrid?

See, e.g., the necessary `-O4` flag with `mpxlf_r` on IBM Power6 systems
Three problems:

- **the application problem:**
  - one must separate application into:
    - **code that can run before the halo data is received**
    - **code that needs halo data**
  
  ➔ very hard to do !!!

- **the thread-rank problem:**
  - comm. / comp. via thread-rank
  - cannot use work-sharing directives
  
  ➔ loss of major OpenMP support (see next slide)

- **the load balancing problem**

```c
if (my_thread_rank < 1) {
    MPI_Send/Recv....
} else {
    my_range = (high-low-1) / (num_threads-1) + 1;
    my_low = low + (my_thread_rank)*my_range;
    my_high=high+ (my_thread_rank+1)*my_range;
    my_high = max(high, my_high)
    for (i=my_low; i<my_high; i++) {
        ....
    }
}
```
Overlapping Communication and Computation

MPI communication by one or a few threads while other threads are computing

Subteams

- Important proposal for OpenMP 3.x or OpenMP 4.x


```c
#pragma omp parallel
{
#pragma omp single onthreads(0)
    {
        MPI_Send/Recv…
    }
#pragma omp for onthreads(1:omp_get_numthreads()-1)
    for (……)
        { /* work without halo information */
            /* barrier at the end is only inside of the subteam */
        }
    ...
#pragma omp barrier
#pragma omp for
    for (……)
        { /* work based on halo information */
        }
} /*end omp parallel */
```
Parallel Programming Models on Hybrid Platforms

- **pure MPI**
  - one MPI process on each core

- **hybrid MPI+OpenMP**
  - MPI: inter-node communication
  - OpenMP: inside of each SMP node

- **OpenMP only**
  - distributed virtual shared memory

- **No overlap of Comm. + Comp.**
  - MPI only outside of parallel regions of the numerical application code

- **Overlapping Comm. + Comp.**
  - MPI communication by one or a few threads while other threads are computing

- **Master only**
  - MPI only outside of parallel regions

- **Multiple/only**
  - appl. threads inside of MPI

- **Funneled**
  - MPI only on master-thread

- **Multiple**
  - more than one thread may communicate

- **Funneled & Reserved**
  - reserved thread for communication

- **Funneled with Full Load Balancing**

- **Multiple & Reserved**
  - reserved threads for communication

- **Multiple with Full Load Balancing**

Different strategies to simplify the load balancing
Experiment: Matrix-vector-multiply (MVM)

- Jacobi-Davidson-Solver on IBM SP Power3 nodes with 16 CPUs per node
  - funneled & reserved is always faster in this experiment
  - Reason:
    Memory bandwidth is already saturated by 15 CPUs, see inset
  - Inset: Speedup on 1 SMP node using different number of threads

NEW OpenMP Tasking Model gives a new way to achieve more parallelism form hybrid computation.


Slides, courtesy of Alice Koniges, NERSC, LBNL
Case study: Communication and Computation in Gyrokinetic Tokamak Simulation (GTS) shift routine

Work on particle array (packing for sending, reordering, adding after sending) can be overlapped with data independent MPI communication using OpenMP tasks.
Overlapping can be achieved with OpenMP tasks (1st part)

- **Overlap**: Master thread encounters (!$omp master) tasking statements and creates work for the thread team for deferred execution. MPI Allreduce call is immediately executed.
- MPI implementation has to support at least MPI_THREAD_FUNNELED
- Subdividing tasks into smaller chunks to allow better load balancing and scalability among threads.

Slides, courtesy of Alice Koniges, NERSC, LBNL
Overlapping can be achieved with OpenMP tasks (2nd part)

Particle reordering of remaining particles (above) and adding sent particles into array (right) & sending or receiving of shifted particles can be independently executed.

Overlapping particle reordering

Overlapping remaining MPI_Sendrecv

Slides, courtesy of Alice Koniges, NERSC, LBNL
OpenMP tasking version outperforms original shifter, especially in larger poloidal domains

- Performance breakdown of GTS shifter routine using 4 OpenMP threads per MPI process with varying domain decomposition and particles per cell on Franklin Cray XT4.
- MPI communication in the shift phase uses a toroidal MPI communicator (constantly 128).
- Large performance differences in the 256 MPI run compared to 2048 MPI run!
- Speed-Up is expected to be higher on larger GTS runs with hundreds of thousands CPUs since MPI communication is more expensive.
OpenMP/DSM

- Distributed shared memory (DSM) //
- Distributed virtual shared memory (DVSM) //
- Shared virtual memory (SVM)

- Principles
  - emulates a shared memory
  - on distributed memory hardware

- Implementations
  - e.g., Intel® Cluster OpenMP
Basic idea:

• Between OpenMP barriers, data exchange is not necessary, i.e., visibility of data modifications to other threads only after synchronization.

• When a page of sharable memory is not up-to-date, it becomes **protected**.

• Any access then faults (SIGSEGV) into Cluster OpenMP runtime library, which requests info from remote nodes and updates the page.

• Protection is removed from page.

• Instruction causing the fault is re-started, this time successfully accessing the data.
Comparison:

MPI based parallelization ↔ DSM

- MPI based:
  - Potential of boundary exchange between two domains in one large message
    → Dominated by bandwidth of the network

- DSM based (e.g. Intel® Cluster OpenMP):
  - Additional latency based overhead in each barrier
    → May be marginal
  - Communication of updated data of pages
    → Not all of this data may be needed
    → i.e., too much data is transferred
    → Packages may be too small
    → Significant latency
  - Communication not oriented on boundaries of a domain decomposition
    → Probably more data must be transferred than necessary

by rule of thumb:
Communication may be 10 times slower than with MPI
Comparing results with heat example

- Normal OpenMP on shared memory (ccNUMA) NEC TX-7

![Graph comparing performance of heat_x.c and heatc2_x.c with OpenMP on NEC TX-7](image)
Heat example: Cluster OpenMP Efficiency

- Cluster OpenMP on a Dual-Xeon cluster

Efficiency only with small communication foot-print

Up to 3 CPUs with 3000x3000

Terrible with non-default schedule

No speedup with 1000x1000

Second CPU only usable in small cases
Back to the mixed model – an Example

- Topology-problem solved: Only horizontal inter-node comm.
- Still intra-node communication
- Several threads per SMP node are communicating in parallel: network saturation is possible
- Additional OpenMP overhead
- With Masteronly style: 75% of the threads sleep while master thread communicates
- With Overlapping Comm. & Comp.: Master thread should be reserved for communication only partially – otherwise too expensive
- MPI library must support
  - Multiple threads
  - Two fabrics (shmemp + internode)
No silver bullet

- The analyzed programming models do not fit on hybrid architectures
  - whether drawbacks are minor or major
    - depends on applications’ needs
  - But there are major opportunities → next section

- In the NPB-MZ case-studies
  - We tried to use optimal parallel environment
    - for pure MPI
    - for hybrid MPI+OpenMP
  - i.e., the developers of the MZ codes and we tried to minimize the mismatch problems
→ the opportunities in next section dominated the comparisons
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Nested Parallelism

- Example NPB: BT-MZ (Block tridiagonal simulated CFD application)
  - Outer loop:
    - limited number of zones \(\Rightarrow\) limited parallelism
    - zones with different workload \(\Rightarrow\) speedup \(<\frac{\text{Sum of workload of all zones}}{\text{Max workload of a zone}}\)
  - Inner loop:
    - OpenMP parallelized (static schedule)
    - Not suitable for distributed memory parallelization

- Principles:
  - Limited parallelism on outer level
  - Additional inner level of parallelism
  - Inner level not suitable for MPI
  - Inner level may be suitable for static OpenMP worksharing
Load-Balancing
(on same or different level of parallelism)

• OpenMP enables
  – Cheap *dynamic* and *guided* load-balancing
  – Just a parallelization option (clause on omp for / do directive)
  – Without additional software effort
  – Without explicit data movement

• On MPI level
  – *Dynamic load balancing* requires moving of parts of the data structure through the network
  – Significant runtime overhead
  – Complicated software / therefore not implemented

• MPI & OpenMP
  – Simple static load-balancing on MPI level, dynamic or guided on OpenMP level

\{ medium quality
  cheap implementation \}
Memory consumption

- Shared nothing
  - Heroic theory
  - In practice: Some data is duplicated

- **MPI & OpenMP**
  With n threads per MPI process:
  - Duplicated data may be reduced by factor n
Using more OpenMP threads could reduce the memory usage substantially, up to five times on Hopper Cray XT5 (eight-core nodes).

Memory consumption (continued)

• Future:
  With 100+ cores per chip the memory per core is limited.
  – Data reduction through usage of shared memory may be a key issue
  – Domain decomposition on each hardware level
    • **Maximizes**
      – Data locality
      – Cache reuse
    • **Minimizes**
      – ccNUMA accesses
      – Message passing
  – No halos between domains inside of SMP node
    • **Minimizes**
      – Memory consumption
How many threads per MPI process?

- SMP node = with **m sockets** and **n cores/socket**
- How many threads (i.e., cores) per MPI process?
  - Too many threads per MPI process
    - overlapping of MPI and computation may be necessary,
    - some NICs unused?
  - Too few threads
    - too much memory consumption (see previous slides)
- Optimum
  - somewhere between 1 and m x n threads per MPI process,
  - Typically:
    - **Optimum** = n, i.e., 1 MPI process per socket
    - **Sometimes** = n/2, i.e., 2 MPI processes per socket
    - **Seldom** = 2n, i.e., each MPI process on 2 sockets
Opportunities, if MPI speedup is limited due to algorithmic problems

- Algorithmic opportunities due to larger physical domains inside of each MPI process
  - If multigrid algorithm only inside of MPI processes
  - If separate preconditioning inside of MPI nodes and between MPI nodes
  - If MPI domain decomposition is based on physical zones
To overcome MPI scaling problems

- Reduced number of MPI messages, reduced aggregated message size compared to pure MPI
- MPI has a few scaling problems
  - Handling of more than 10,000 MPI processes
  - Irregular Collectives: MPI_..v(), e.g. MPI_Gatherv()
    - Scaling applications should not use MPI_..v() routines
  - MPI-2.1 Graph topology (MPI_Graph_create)
    - MPI-2.2 MPI_Dist_graph_create_adjacent
  - Creation of sub-communicators with MPI_Comm_create
    - MPI-2.2 introduces a new scaling meaning of MPI_Comm_create
- Hybrid programming reduces all these problems (due to a smaller number of processes)
Summary: Opportunities of hybrid parallelization (MPI & OpenMP)

- Nested Parallelism
  - Outer loop with MPI / inner loop with OpenMP

- Load-Balancing
  - Using OpenMP *dynamic* and *guided* worksharing

- Memory consumption
  - Significantly reduction of replicated data on MPI level

- Opportunities, if MPI speedup is limited due to algorithmic problem
  - Significantly reduced number of MPI processes

- Reduced MPI scaling problems
  - Significantly reduced number of MPI processes
Outline

- Introduction / Motivation
- Programming models on clusters of SMP nodes
- Case Studies / Benchmark results
- Mismatch Problems
- Opportunities:
  Application categories that can benefit from hybrid parallelization

- **Thread-safety quality of MPI libraries**

- Other options on clusters of SMP nodes
- Summary
MPI rules with OpenMP / Automatic SMP-parallelization

- Special MPI-2 Init for multi-threaded MPI processes:

```c
int MPI_Init_thread( int * argc, char ** argv[],
                     int thread_level_required,
                     int * thread_level_provided);
int MPI_Query_thread( int * thread_level_provided);
int MPI_Is_main_thread(int * flag);
```

- REQUIRED values (increasing order):
  - `MPI_THREAD_SINGLE`: Only one thread will execute
  - `THREAD_MASTERONLY`: MPI processes may be multi-threaded, but only master thread will make MPI-calls AND only while other threads are sleeping
  - `MPI_THREAD_FUNNELED`: Only master thread will make MPI-calls
  - `MPI_THREAD_SERIALIZED`: Multiple threads may make MPI-calls, but only one at a time
  - `MPI_THREAD_MULTIPLE`: Multiple threads may call MPI, with no restrictions

- returned `provided` may be less than REQUIRED by the application
Calling MPI inside of OMP MASTER

• Inside of a parallel region, with “OMP MASTER”

• Requires MPI_THREAD_FUNNELED, i.e., only master thread will make MPI-calls

• **Caution:** There isn’t any synchronization with “OMP MASTER”! Therefore, “OMP BARRIER” normally necessary to guarantee, that data or buffer space from/for other threads is available before/after the MPI call!

  ```
  !$OMP BARRIER #pragma omp barrier
  !$OMP MASTER #pragma omp master
  call MPI_Xxx(…)
  !$OMP END MASTER
  !$OMP BARRIER #pragma omp barrier
  ```

• But this implies that all other threads are sleeping!
• The additional barrier implies also the necessary cache flush!
... the barrier is necessary – example with MPI_Recv

```c
#pragma omp parallel
{
    #pragma omp for nowait
    for (i=0; i<1000; i++)
        a[i] = buf[i];

    #pragma omp barrier
    #pragma omp master
    MPI_Recv(buf,...);
    #pragma omp barrier

    #pragma omp for nowait
    for (i=0; i<1000; i++)
        c[i] = buf[i];

} /* omp end parallel */
```
Thread support in MPI libraries

- The following MPI libraries offer thread support:

<table>
<thead>
<tr>
<th>Implementation</th>
<th>Thread support level</th>
</tr>
</thead>
<tbody>
<tr>
<td>MPIch-1.2.7p1</td>
<td>Always announces MPI_THREAD_FUNNELED.</td>
</tr>
<tr>
<td>MPIch2-1.0.8</td>
<td>ch3:sock supports MPI_THREAD_MULTIPLE</td>
</tr>
<tr>
<td></td>
<td>ch:nemesis has “Initial Thread-support”</td>
</tr>
<tr>
<td></td>
<td>ch3:nemesis (default) has MPI_THREAD_MULTIPLE</td>
</tr>
<tr>
<td>MPIch2-1.1.0a2</td>
<td>Full MPI_THREAD_MULTIPLE</td>
</tr>
<tr>
<td>Intel MPI 3.1</td>
<td>MPI_THREAD_FUNNELED</td>
</tr>
<tr>
<td>SciCortex MPI</td>
<td>Full MPI_THREAD_MULTIPLE (with libmtmpi)</td>
</tr>
<tr>
<td>HP MPI-2.2.7</td>
<td>Not thread-safe?</td>
</tr>
<tr>
<td>SGI MPT-1.14</td>
<td>Full MPI_THREAD_MULTIPLE</td>
</tr>
<tr>
<td>IBM MPI</td>
<td>MPI_THREAD_SERIALIZED</td>
</tr>
<tr>
<td>Nec MPI/SX</td>
<td></td>
</tr>
</tbody>
</table>

- Testsuites for thread-safety may still discover bugs in the MPI libraries
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- **Other options on clusters of SMP nodes**

- Summary
Pure MPI – multi-core aware

- Hierarchical domain decomposition (or distribution of Cartesian arrays)

Domain decomposition:
1 sub-domain / SMP node

Further partitioning:
1 sub-domain / socket

1 / core

Cache optimization:
Blocking inside of each core, block size relates to cache size. 1-3 cache levels.

Example on 10 nodes, each with 4 sockets, each with 6 cores.
How to achieve a hierarchical domain decomposition (DD)?

• Cartesian grids:
  – Several levels of subdivide
  – Ranking of MPI_COMM_WORLD – two choices:
    a) Sequential ranks through original data structure
       + locating these ranks correctly on the hardware
       ➢ can be achieved with one-level DD on finest grid
         + special startup (mpiexec) with optimized rank-mapping
    b) Sequential ranks in comm_cart (from MPI_CART_CREATE)
       ➢ requires optimized MPI_CART_CREATE,
         or special startup (mpiexec) with optimized rank-mapping
    c) Sequential ranks in MPI_COMM_WORLD
       + additional communicator with sequential ranks in the data structure
       + self-written and optimized rank mapping.

• Unstructured grids:
  ➔ next slide
How to achieve a hierarchical domain decomposition (DD)?

- Unstructured grids:
  - Multi-level DD:
    - Top-down: Several levels of (Par)Metis
    - Bottom-up: Low level DD + higher level recombination
  - Single-level DD (finest level)
    - Analysis of the communication pattern in a first run (with only a few iterations)
    - Optimized rank mapping to the hardware before production run
    - E.g., with CrayPAT + CrayApprentice
Top-down – several levels of (Par)Metis

Steps:
- Load-balancing (e.g., with ParMetis) on outer level, i.e., between all SMP nodes
- Independent (Par)Metis inside of each node
- Metis inside of each socket

⚠️ Subdivide does not care on balancing of the outer boundary

⚠️ processes can get a lot of neighbors with inter-node communication

⚠️ unbalanced communication
Bottom-up –
Multi-level DD through recombination

1. Core-level DD: partitioning of application’s data grid
2. Socket-level DD: recombining of core-domains
3. SMP node level DD: recombining of socket-domains

• Problem:
  Recombination must not calculate patches that are smaller or larger than the average

• In this example the load-balancer must combine always
  ▪ 6 cores, and
  ▪ 4 sockets

• Advantage:
  Communication is balanced!
Profiling solution

• First run with profiling
  – Analysis of the communication pattern

• Optimization step
  – Calculation of an optimal mapping of ranks in MPI_COMM_WORLD to the hardware grid (physical cores / sockets / SMP nodes)

• Restart of the application with this optimized locating of the ranks on the hardware grid

• Example: CrayPat and CrayApprentice
Scalability of MPI to hundreds of thousands ...

Weak scalability of pure MPI

- As long as the application does not use
  - `MPI_ALLTOALL`
  - `MPI_<collectives>V` (i.e., with length arrays)
  and application
  - distributes all data arrays
  one can expect:
- Significant, but still scalable memory overhead for halo cells.
- MPI library is internally scalable:
  - E.g., mapping ranks \(\rightarrow\) hardware grid
    - Centralized storing in shared memory (OS level)
    - In each MPI process, only used neighbor ranks are stored (cached) in process-local memory.
  - Tree based algorithm with \(O(\log N)\)
    - From 1000 to 1000,000 process \(O(\log N)\) only doubles!

The vendors will (or must) deliver scalable MPI libraries for their largest systems!
Remarks on Cache Optimization

- **After** all parallelization domain decompositions (DD, up to 3 levels) are done:
- Additional DD into data blocks
  - that fit to 2nd or 3rd level cache.
  - It is done inside of each MPI process (on each core).
  - Outer loops over these blocks
  - Inner loops inside of a block
- Cartesian example: 3-dim loop is split into

```fortran
  do i_block=1,ni,stride_i
    do j_block=1,nj,stride_j
      do k_block=1,nk,stride_k
        do i=i_block,min(i_block+stride_i-1, ni)
          do j=j_block,min(j_block+stride_j-1, nj)
            do k=k_block,min(k_block+stride_k-1, nk)
              a(i,j,k) = f( b(i±0,1,2, j±0,1,2, k±0,1,2) )
              ... ... ... end do
            end do
          end do
        end do
      end do
    end do
  end do
```

Access to 13-point stencil
Remarks on Cost-Benefit Calculation

Costs
• for optimization effort
  – e.g., additional OpenMP parallelization
  – e.g., 3 person month x 5,000 € = 15,000 € (full costs)

Benefit
• from reduced CPU utilization
  – e.g., Example 1:
    100,000 € hardware costs of the cluster
    x 20% used by this application over whole lifetime of the cluster
    x 7% performance win through the optimization
    = 1,400 € → total loss = 13,600 €
  – e.g., Example 2:
    10 Mio € system x 5% used x 8% performance win
    = 40,000 € → total win = 25,000 €
Remarks on MPI and PGAS (UPC & CAF)

- Parallelization always means expressing locality.

- If the application has no locality,
  - Then the parallelization needs not to model locality

  \[\text{UPC with its round robin data distribution may fit}\]

- If the application has locality,
  - then it must be expressed in the parallelization

- Coarray Fortran (CAF) expresses data locality explicitly through “co-dimension”:
  - \[A(17,15)[3]\]
    = element \(A(17,13)\) in the distributed array \(A\) in process with rank 3
Remarks on MPI and PGAS (UPC & CAF)

• Future shrinking of memory per core implies
  – Communication time becomes a bottleneck
  → Computation and communication must be overlapped,
    i.e., latency hiding is needed

• With PGAS, halos are not needed.
  – But it is hard for the compiler to access data such early that the
    transfer can be overlapped with enough computation.

• With MPI, typically too large message chunks are transferred.
  – This problem also complicates overlapping.

• Strided transfer is expected to be slower than contiguous transfers
  – Typical packing strategies do not work for PGAS on compiler level
  – Only with MPI, or with explicit application programming with PGAS
Remarks on MPI and PGAS (UPC & CAF)

- Point-to-point neighbor communication
  - PGAS or MPI nonblocking may fit
    if message size makes sense for overlapping.

- Collective communication
  - Library routines are best optimized
  - Non-blocking collectives (comes with MPI-3.0)
    versus calling MPI from additional communication thread
  - Only blocking collectives in PGAS library?
Remarks on MPI and PGAS (UPC & CAF)

- For extreme HPC (many nodes \times many cores)
  - Most parallelization may still use MPI
  - Parts are optimized with PGAS, e.g., for better latency hiding
  - PGAS efficiency is less portable than MPI
  - `#ifdef ... PGAS`
  - Requires mixed programming PGAS & MPI
    \[ \rightarrow \text{will be addressed by MPI-3.0} \]
Outline

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• Opportunities:
  Application categories that can benefit from hybrid parallelization
• Thread-safety quality of MPI libraries
• Other options on clusters of SMP nodes

• Summary
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  - HPCMO Program and the Engineer Research and Development Center Major Shared Resource Center, Vicksburg, MS (http://www.erdc.hpc.mil/index)
Summary – the good news

MPI + OpenMP

• Significant opportunity → higher performance on smaller number of threads
• Seen with NPB-MZ examples
  – BT-MZ → strong improvement (as expected)
  – SP-MZ → small improvement (none was expected)
• Usable on higher number of cores
• Advantages
  – Load balancing
  – Memory consumption
  – Two levels of parallelism
    • Outer → distributed memory → halo data transfer → MPI
    • Inner → shared memory → ease of SMP parallelization → OpenMP
• You can do it → “How To”
Summary – the bad news

MPI+OpenMP: There is a huge amount of pitfalls

• Pitfalls of MPI
• Pitfalls of OpenMP
  – On ccNUMA → e.g., first touch
  – Pinning of threads on cores
• Pitfalls through combination of MPI & OpenMP
  – E.g., topology and mapping problems
  – Many mismatch problems
• Tools are available 😊
  – It is not easier than analyzing pure MPI programs 😞
• Most hybrid programs → Masteronly style
• Overlapping communication and computation with several threads
  – Requires thread-safety quality of MPI library
  – Loss of OpenMP worksharing support → using OpenMP tasks as workaround
Summary – good and bad

- Optimization
  - 1 MPI process mismatch
  - per core ......................................................
     ^– somewhere between
  - 1 MPI process
     per SMP node
     may be the optimum

- 😊 Efficiency of MPI+OpenMP is not for free:
  - The efficiency strongly depends on
  - 😏 the amount of work in the source code development
Summary – Alternatives

Pure MPI
+ Ease of use
  – Topology and mapping problems may need to be solved
    (depends on loss of efficiency with these problems)
  – Number of cores may be more limited than with MPI+OpenMP
+ Good candidate for perfectly load-balanced applications

Pure OpenMP
+ Ease of use
  – Limited to problems with tiny communication footprint
  – source code modifications are necessary
    (Variables that are used with “shared” data scope
     must be allocated as “sharable”)
± (Only) for the appropriate application a suitable tool
Summary

- This tutorial tried to
  - help to negotiate obstacles with hybrid parallelization,
  - give hints for the design of a hybrid parallelization,
  - and technical hints for the implementation → “How To”,
  - show tools if the application does not work as designed.

- This tutorial was not an introduction into other parallelization models:
  - Partitioned Global Address Space (PGAS) languages
    (Unified Parallel C (UPC), Co-array Fortran (CAF), Chapel, Fortress, Titanium, and X10).
  - High Performance Fortran (HPF)
  → Many rocks in the cluster-of-SMP-sea do not vanish into thin air by using new parallelization models
  → Area of interesting research in next years
Conclusions

• Future hardware will be more complicated
  – Heterogeneous → GPU, FPGA, ...
  – ccNUMA quality may be lost on cluster nodes
  – ....
• High-end programming → more complex
• Medium number of cores → more simple
  (if `#cores / SMP-node` will not shrink)
• MPI+OpenMP → work horse on large systems
• Pure MPI → still on smaller cluster
• OpenMP → on large ccNUMA nodes
  (not ClusterOpenMP)

Thank you for your interest

Q & A

Please fill in the feedback sheet – Thank you
Appendix

- Author
- References (with direct relation to the content of this tutorial)
- Further references
Rolf Rabenseifner

Dr. Rolf Rabenseifner studied mathematics and physics at the University of Stuttgart. Since 1984, he has worked at the High-Performance Computing-Center Stuttgart (HLRS). He led the projects DFN-RPC, a remote procedure call tool, and MPI-GLUE, the first metacomputing MPI combining different vendor's MPIs without losing the full MPI interface. In his dissertation, he developed a controlled logical clock as global time for trace-based profiling of parallel and distributed applications. Since 1996, he has been a member of the MPI-2 Forum and since Dec. 2007, he is in the steering committee of the MPI-3 Forum. From January to April 1999, he was an invited researcher at the Center for High-Performance Computing at Dresden University of Technology. Currently, he is head of Parallel Computing - Training and Application Services at HLRS. He is involved in MPI profiling and benchmarking, e.g., in the HPC Challenge Benchmark Suite. In recent projects, he studied parallel I/O, parallel programming models for clusters of SMP nodes, and optimization of MPI collective routines. In workshops and summer schools, he teaches parallel programming models in many universities and labs in Germany.
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